

### HARVARD COLLEGE OBSERVATORY

60 Garden Street, Cambridge, MA 02138

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Title: Measurements of Ultraviolet Absorption Cross Sections of

NO<sub>2</sub> at Various Pressures and Temperatures

Principal Investigator: W. H. Parkinson

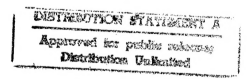
Co-Investigator: Kouichi Yoshino

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60 Garden Street, MS-50 Cambridge, MA 02138

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## Abstract

Laboratory measurements of the absorption cross sections of  $NO_2$  at the temperature 298.5 K have been made throughout the wavelength region 360 nm to 470 nm. The equilibrium constant  $K_p$  has been experimentally obtained as 150 hPa, from which the column density of pure  $NO_2$  is obtained. The cross sections almost completely agree with the recent results of Mérienne  $et\ al.$  [J. Atmos. Chem. 20, 281-297 (1995)]. The cross sections at the higher resolution of 0.14 cm<sup>-1</sup> have been obtained in the wavelength region where the spectra have fine structure.

W. H. Parkinson

Kouichi Yoshino

### 1 Introduction

NO<sub>2</sub> plays a critical role in stratospheric ozone chemistry. The nitrogen oxides, NO and NO<sub>2</sub>, are introduced into the atmosphere by the reaction of N<sub>2</sub>O, which is naturally produced by bacterial action on organic material in the ground. NO<sub>2</sub>, produced at the surface, diffuses upwards into the atmosphere, where, in collision with atomic oxygen, NO is formed. NO in reaction with ozone is rapidly converted to nitrogen dioxide. The photodissociation of NO<sub>2</sub> and its increasing presence as an atmospheric pollutant, combine to make it an important participant in the chemistry of the lower atmosphere.

Knowledge of the absorption cross section of NO<sub>2</sub> is requires in order to interpret measurements, and for calculation of the rate coefficient for NO<sub>2</sub> photodissociation in the atmosphere.

The electronic absorption spectrum of NO<sub>2</sub> is known to extend from 320 to 1000 nm. The spectrum consists of a very large number of bands, and there is no obvious regularity either in the vibrational structure or in the rotational structure of the individual bands. The analysis of these bands has been a long-standing spectroscopic problem which is still largely unsolved. Douglas and Huber [1965] investigated the six absorption bands of <sup>14</sup>NO<sub>2</sub> and <sup>15</sup>NO<sub>2</sub> in the region 370 -460 nm, and assigned them to  $K_a = 0 \leftarrow 1$  subbands of the electronic transition  $^2B_1 \leftarrow^2 A_1$ .

In preparation for photoelectric measurements of the absolute ultra violet absorption cross section of NO<sub>2</sub>, we borrowed the high resolution, photographic plates of NO<sub>2</sub> absorption by Douglas and Huber [1965] and studied what wavelength resolution was necessary for the cross section measurements of NO<sub>2</sub>. We had planned to use these spectra for wavelengths calibration of our cross section measurements, but abandoned the plan because of the lack of isolated sharp lines in the bands.

# 2 Experimental Procedure

We used the 6.65-m, normal incidence, vacuum spectrometer in the first order of a 1200  $l/\mathrm{mm}$  grating to obtain photoabsorption cross section measurements of NO<sub>2</sub>. The grating was blazed at 5500 Å, and provided an estimated instrumental width of 0.0025 nm with entrance and exit slit widths of 0.010 mm. The stainless steel absorption cell was connected directly to the entrance slit assembly of the spectrometer and was closed with two fused silica windows (Fig. 1). The path length was 42.3 cm. The continuous background for the photoabsorption measurements was provided by a tungsten lamp operated on 12 V ac. The NO<sub>2</sub> gas (Matheson, 99.5%) was purified

by the method of fractional distillation, and was kept in a liquid-nitrogen cooled trap. The NO<sub>2</sub> was warmed up slowly and introduced into the absorption cell. NO<sub>2</sub> pressures of 0.5, 1.0, 1.5, 2.0, 2.5, and 3.0 Torr were used and measured with a capacitance manometer (MKS Baratron, 10 Torr).

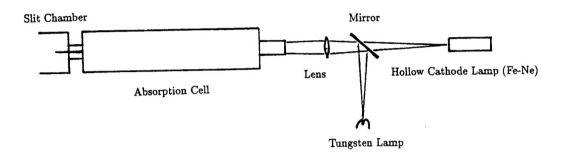


Figure 1: Experimental arrangement of absorption cell, background continuum source, and hollow cathode lamp.

We divided the spectral region (360 nm - 470 nm) into sixteen sections of about 7.5 nm extent. The background continuum level was established by scanning the empty cell before and after each photoabsorption measurements, and comparing with the background intensity levels without NO<sub>2</sub> in the cell. We obtained data points every 0.0015 nm at a resolution of 0.0040 nm. The cross section measurements for the most structured region were also obtained with a resolution of 0.0025 nm for every 0.0003 nm step. In each scan range, two extra scans were obtained; the back ground scan without any NO<sub>2</sub> in the cell and the scan of the hollow cathode lamp (Fe/Ne, 20 mA) for wavelength calibration. Some times we also took a scan of both the main NO<sub>2</sub> spectrum and the hollow cathode spectrum to avoid any systematic shift in wavelength. The wavelengths of Fe I, Fe II, and Ne I lines are well known (± 0.001 Å). The uncertainty of the wavelength calibration depended on the scan step, which was 0.015 Å and for the high resolution scan, 0.003 Å.

Most of the uncertainty in cross section measurements with a single beam spectrometer is in the estimation of the background intensity levels since they vary with time and wavelength during photoabsorption scan. It is also difficult to keep the column density of NO<sub>2</sub> constant in the absorption cell during the scan. To overcome these problems, we measured the cross section at the very beginning of every scan when the NO<sub>2</sub> gas was fresh, and the background, I<sub>0</sub>, was well known. We obtained these absolute points for sixteen wavelengths and six pressures. The observed cross

sections are a linear function of pressure, and therefore a linear extrapolation to zero pressure provides the cross sections of pure NO<sub>2</sub>. The resulting cross sections at zero pressure,  $\sigma_0$ , are given in Table I along with slopes. The cross sections ( $\sigma_0$ ) can be used as the absolute cross sections. The ratio of the observed to absolute cross sections gives the fraction of NO<sub>2</sub> density at that pressure, and these are given in Table 2 for fourteen runs. The averaged fractions are given in the same table with the uncertainties ( $1\sigma$  and  $\sigma/\sqrt{n}$ ), and are also plotted in Fig. 2. The equilibrium constants  $K_p$  are given by

 $K_p = \frac{p^2(NO_2)}{p(N_2O_4)}. (1)$ 

The dotted curves in the figure are the fractions vs. pressures calculated from values of  $K_p$  with the curves in order from the top,  $K_p$  of 160, 150, and 140hPa. Our observed fractions agree very well with the fraction calculated with  $K_p = 150$ . Hurtmans et al. [1993] determined the  $K_p$  value by fitting the literature values with a fourth degree polynomial expansion in the temperature. According to Roscoe and Hind [1993], the error in the literature values of the equilibrium constants is about 5%, but we could estimate the constants with an uncertainty of better than 2%.

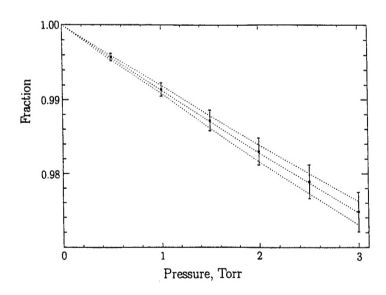


Figure 2: The observed fractions of NO<sub>2</sub> plotted against the total pressure. The dotted curves are calculated fractions of NO<sub>2</sub> with  $K_p = 160$ , 150, and 140 (in the order from the top). The observation agree with  $K_p = 150$ , which is expected for a temperature of 298.54 K.

Table 1. Cross Sections of  $NO_2$  at 295 K and pressure dependencies.

Run	Wavenumber cm <sup>-1</sup>	Wavelength nm(air)	$\frac{\sigma_0}{10^{-19} \text{cm}^2}$	Slope 10 <sup>-19</sup> cm <sup>2</sup> /Torr
				10 "cm"/ for
R35	27824.74	359.2899	$4.56 \pm 0.20$	0.0306
R36	27242.44	366.9698	$5.29 \pm 0.18$	-0.0206
R37	26685.10	374.6345	$5.19 \pm 0.05$	-0.0381
R38	26137.12	382.4891	$5.44 \pm 0.05$	-0.0285
R39	25599.86	390.5165	$6.17 \pm 0.06$	-0.0619
R40	25164.98	397.2652	$5.56 \pm 0.03$	-0.0706
R41	24744.51	404.0159	$6.38 \pm 0.05$	-0.0783
R42	24346.00	410.6289	$6.82 \pm 0.02$	-0.0283
R43	23953.98	417.3496	$5.54 \pm 0.03$	-0.0480
R44	23576.06	424.0398	$6.34 \pm 0.08$	-0.1203
R45	23218.81	430.5641	$6.10 \pm 0.04$	-0.0609
R46	22864.07	437.2446	$4.31 \pm 0.03$	-0.0614
R47	22521.29	443.8997	$4.44 \pm 0.02$	-0.0548
R48	22244.90	449.4152	$4.31 \pm 0.08$	-0.0376
R49	21873.73	457.0414	$3.83 \pm 0.02$	-0.0358
R50	21533.08	464.2717	$4.15 \pm 0.03$	-0.0102

Table 2. Fractions of NO<sub>2</sub> density versus pressure

P(NO <sub>2</sub> ), Torr	0.5	1.0	1.5	2.0	2.5	3.0
R36	0.9981	0.9961	0.9942	0.9923	0.9904	0.9885
R37	0.9963	0.9927	0.9891	0.9855	0.9820	0.9784
R38	0.9974	0.9948	0.9922	0.9896	0.9871	0.9845
R39	0.9950	0.9901	0.9852	0.9803	0.9755	0.9708
R40	0.9937	0.9875	0.9813	0.9752	0.9692	0.9633
R41	0.9939	0.9879	0.9819	0.9760	0.9702	0.9645
R42	0.9979	0.9959	0.9938	0.9918	0.9897	0.9877
R43	0.9957	0.9914	0.9872	0.9830	0.9788	0.9747
R45	0.9950	0.9901	0.9852	0.9804	0.9756	0.9709
R46	0.9929	0.9860	0.9791	0.9723	0.9656	0.9590
R47	0.9939	0.9878	0.9818	0.9759	0.9701	0.9643
R48	0.9957	0.9913	0.9871	0.9828	0.9786	0.9745
R49	0.9954	0.9908	0.9862	0.9817	0.9772	0.9728
R50	0.9988	0.9975	0.9963	0.9951	0.9939	0.9927
Average	0.9957	0.9914	0.9872	0.9830	0.9789	0.9748
$\sigma$	0.0017	0.0034	0.0051	0.0068	0.0085	0.0101
$\sigma/\sqrt{n}$	0.0005	0.0009	0.0014	0.0018	0.0023	0.0027

## 3 The cross sections of $NO_2$

The cross sections of every scan were obtained by the Beer-Lambert law assuming that the observed pressure represents the pure NO<sub>2</sub> concentration, and later were calibrated to the true cross sections by using the fractions of NO<sub>2</sub> density shown in Table 2. The cross sections at the beginning of the scan presented in Table 1 are the absolute cross sections because the column densities are obtained with a fresh NO<sub>2</sub> sample. The cross sections at the end of scan are calibrated with the beginning value of the following scan of the next wavelength region with the same pressure. All cross sections obtained with different pressures are put together to reduce the final cross sections of the each sections. However data with optical depths higher than 2.0 and lower than 0.30 are omitted from summations. Our final cross sections are presented in Fig. 3 for the wavelength region 360 to 470 nm. These cross sections agree so well with the recent measurements by Mérienne et al. [1995], that there are no measurable differences on this scale.

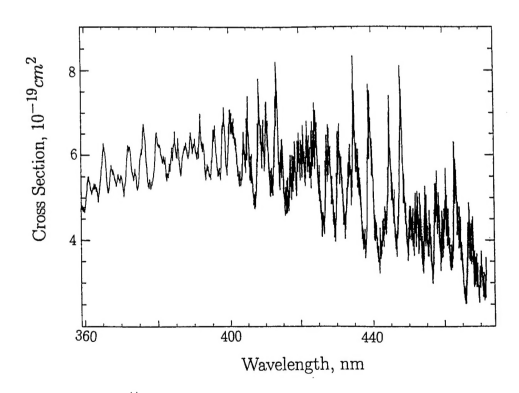


Figure 3: The absorption cross sections of  $NO_2$  at 298.5 K in the wavelength region 360 to 470 nm (air).

Table 3. Wavelength Region of High Resolution Cross Sections

File	Wavelength	Region, nm
	begining	end
411	404.8	407.2
412/3	407.7	410.8
421/2	412.4	415.5
431	423.2	424.8
441	426.9	428.6
451	430.1	431.5
452/3	433.4	436.8
461/2	438.3	441.4
471	444.0	445.7
472	447.3	449.1
481	454.3	455.7
482	451.7	453.1
491	462.7	464.1

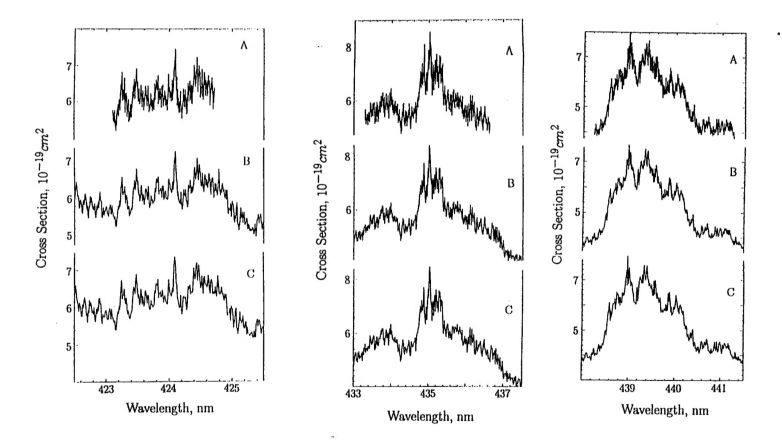


Figure 4: (A): The absorption cross sections of  $NO_2$  at the high resolution of 0.14 cm<sup>-1</sup>. (B): the cross sections at the resolution of 0.28 cm<sup>-1</sup>. (C) The cross sections of Mérienne *et al.* [1995] at the resolution of 0.50 cm<sup>-1</sup>.

The cross section measurements at high resolution covered several wavelength region, where the fine band structures are observed. The high resolution scan used the instrumental width of 0.0025 nm, and data points were taken every 0.003 nm step. The wavelength region for the high resolution scan are tabulated in Table 3. A few examples of the high resolution scan are presented in Fig. 4 through 6 for wavelength region of around 424, 435, and 440 nm. The figures also show our regular scans and those of Mérienne et al. [1995] for comparison. Except in the details, there are almost no differences among the three scans.

The three recent independent measurements of NO<sub>2</sub> cross sections [Mérienne et al., 1995; Vandacle et al., 1996; and ours] in the wavelength region 370-470 nm agree very well. The three groups are planing to combine all these recent data to obtain the recommended cross sections of NO<sub>2</sub> for the convenience of application in atmospheric studies. The cross sections reported here are available at a wavelength interval of 0.005 nm on the World Wide Web. The URL is http://cfa-www.harvard.edu/amp/data/amdata.html.

The cross sections of NO<sub>2</sub> mixed with air and nitrogen gas, were also measured at the same conditions. The values of the cross sections of NO<sub>2</sub> mixed with nitrogen are more or less the same as those of the pure NO<sub>2</sub> gas. However, the cross sections measurements of NO<sub>2</sub> mixed with air lead to very low values compared with those of the pure gas. The re-analysis of the mixed gases composition could not explain why smaller cross sections were observed with NO<sub>2</sub> in the air. We also measured the cross section of NO<sub>2</sub> at 225 K, but it will take more time for the data analysis because the analysis must include consideration of the dimer population.

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#### REFERENCES

A.E. Douglas and K.P. Huber, The Absorption Spectrum of NO<sub>2</sub> in the 3700-4600 Å Region, Can. J. Phys. 43, 74-81 (1965).

D. Hurtmans, M. Herman, and J.V. Auwera, Integrated Band Intensities in

N<sub>2</sub>O<sub>4</sub> in the Infrared Range, J. Quant. Spectrosc. Radiat. Transf. **50**, 595-602 (1993).

M.F. Mérienne, Λ. Jenouvrier, and B. Coquart, The NO<sub>2</sub> Absorption Spectrum. I: Absorption Cross-Sections at Ambient Temperature in the 300-500 nm Region, J. Atmos. Chem. 20, 281-297 (1995).

H.K. Roscoe and A.K. Hind, The Equilibrium Constant of NO<sub>2</sub> with N<sub>2</sub>O<sub>4</sub> and the Temperature Dependence of the Visible Spectrum of NO<sub>2</sub>: A Critical Review and the Implications for Measurements of NO<sub>2</sub> in the Polar Stratosphere, J. Atm. Chem. 16, 257-276 (1993).

A.C. Vandaele, C. Hermans, P.C. Simon, M. Van Roozendael, J.M. Guilmot, M. Carleer, and R. Colin, J. Atm. Chem., (1996). in press.